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# Identification and Procaryotic Expression of the Gene Coding for the Highly Immunogenic 28-Kilodalton Structural Phosphoprotein (pp28) of Human Cytomegalovirus

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Human cytomegalovirus contrins a structural polypeptide that is 28 kilodaltons in apparent molecular size and is reactive in Western blot (immunoblot) analysis with the majority of human sera. The gene coding for this polypeptide was mapped on the genome of human cytomegalovirus strain AD169. A monoclosal antibody specific for the 28-kilodalton polypeptide was used to screen a cDNA library constructed from poly(A)\* RNA of human cytomegalovirus-infected cells in the procuryotic expression vector lambda gt11. Hybridization of cDNA with cosmid and plasmid clones mapped the gene to the Hindill R fragment. The gene was transcribed into a late 1.3-kilobase RNA. The nucleotide sequence of the coding region was determined. Parts of the 28-kilodalton polypeptide were expressed in Escherichis coll as hybrid proteins fused to β-galactosidase. In Western blots these proteins were recognized by human sera. Antibodies raised against the hybrid proteins reacted specifically with the viral antigen in immunoprecipitations and Western blots. In vitro phosphorylated.

Human cytomegalovirus (HCMV), a ubiquitous member of the herpesvirus family, can be associated with a wide spectrum of disease, particularly in immunocompromised persons. The factors that are responsible for this wide spectrum of clinical manifestation of HCMV infections are largely unknown. The route of injection and variation in the strain of infecting virus may influence the development of disease. Host immunological functions clearly play an important role in the control of infection. To more fully understand the importance of the immune response in limiting the severity of HCMV infections, it is necessary to first characterize the virus-encoded targets of this response. Serological diagnosis of active HCMV infection is mainly based on a significant rise in antibody levels detected by a complement-fixation test or an enzyme-linked immunosorbent assay. While highly useful in detecting an immune response to the virus, these assays do not determine the fine specificity of antiviral antibodies since each assay indiscriminately detects antibodies against multiple viral antigens. To give a more precise definition of the humoral immune response during active HCMV infection it is necessary to evaluate titers against defined antigens. HCMV is a highly complex virus encoding approximately 30 to 40 structural proteins and an unknown number of nonstructural polypeptides (for a review, see M. P. Landini and S. Michelson, Prog. Med. Virol., in press). Several laboratories have employed Western blotting (immunoblotting) analysis or immunoprecipitations to identify individual viral polypeptides that are recognized by human immune sera (14, 15, 19; B. Nowak, Ph.D. thesis, University Erlangen-Numberg. 1984). In these studies it was found that immune sera contain various amounts of antibodies against the majority of the structural components of HCMV. The immune response involved 15 t 25 proteins ranging in molecular size from 28 to 200 kilodaltons (kDa) (14, 37). However the intensity and

frequency of the immune reaction was particularly high for four polypeptides of 150, 65, 58, and 28 kDa.:Three of the proteins which elicited this antibody response were characteristics. terized. They corresponded to the basic phosphoprotein (pp150), the major matrix protein (pp65), and the major glycoprotein (ap58). As an initial step in studying the immunogenic potential of various structural polypeptides. started to isolate the genes coding for these highly immuno reactive profeins. This information will be used to synthesize viral antigens via procaryotic or eucaryotic expression systems and to measure antibody levels in a defined system. The mapping and the characterization of the genes coding for pp150, pp65, and gp58 have been reported (9, 17, 24) The approach to the characterization of pp150 and gp58 was th isolation of clones coding for these polypeptides by process yotic expression cloning. A cDNA library from poly(A) KNA, isolated from HFF ceils at late times in infection, w constructed in the vector lambda gt11 (17). In this system the cDNA is inserted into the 3' end of the lacZ gene. Expres sion of the gene results in the synthesis of a fusion protein of β-galactosidase and the respective foreign polypeptide which can be detected by monospecific antisers or monoclo nal antibodies. Here we describe the identification and characterization of the gene coding for the fourth highly immunogenic structural polypeptide of HCMV, the 28-kg polyreptide, which is also shown to be a phosphoprotein We also show that a recombinant Escherichia coll protein containing parts of pp28, can be used to detect HCI antibodies in human sera.

#### MATERIALS AND METHODS

Virus, cell culture, and virion purification procedure. HCMV strain AD169 was propagated in human foreskin fibroblasts by standard methods. Extracellular virus purified from the tissue culture supermatant through a street collectrate gradient (31).

Recombinant plasmids. Plasmids pRR1, pCMS0071

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pCM5009 were described previously (25). Plasmid p271 was generated by inserting the 500-base-pair (bp) EcoRI-Smal fragment of plasmid pHM7 into pEX-2 (29). pHM7 contains the 500-bp KpnI-Smal fragment inserted into M13mp11 (nucleotides 1443 through 1936).

Construction and screening of the cDNA library. The construction and screening of the lambda gt11 library have

been described in detail (9, 17, 36).

Sequence analysis of the viral DNA. Sequencing of lambda gt11 cDNA and viral DNA was performed by the chain termination method (26). The cDNA was subcloned into M13mp11 before sequence analysis. Sequence analysis of HIndIII R fragment was carried out using the M13-chain termination method (26) with the random strategy of Bankier and Barrell (2). The fragment was excised from the plasmid, which was kindly provided by J. Oram and P. Greenaway. The sequence was determined on both strands by the shotgun method, and each base was determined an average of 6.36 times. The DNA sequence was compiled with the DBOTIL program (27) and analyzed with the ANALYSEQ program (28).

Induction of fusion proteins. Fusion proteins of gt11 clones were produced as described previously (36). P271 fusion proteins were produced in *E. coli* pop2136 (29). The cells were grown to a density (A<sub>600</sub>) of 0.2 to 0.3 at 30°C and induced by a quick temperature shift to 42°C. Synthesis of the fusion protein was allowed to continue for 90 min. After that the cells were harvested, lysed in sodium dodecyl sulfate (SDS)-gel sample buffer, and analyzed on SDS-polyscrylamide gel electrophoresis.

Protein gel electrophoresis and Western blot analysis. Polypeptides were denatured in SDS, electrophoretically separated on polyacrylamide gels, and transferred to nitrocellulose as described previously (13, 34).

Production of monospecific anthera against fusion proteins. E. coll extracts containing the fusion proteins were acparated on preparative 8% polyacrylamide gels, stained with Coomassie brilliant blue, and partially destained. The band corresponding to the fusion protein was cut out and frozen in liquid nitrogen. The frozen polyacrylamide gel slices were pulverized in a Dysmembrator (Fa. Braun, Melsungen, Federal Republic of Germany), and the proteins were extracted by repeated incubations with 0.1 M (NH<sub>4</sub>)HCO<sub>2</sub> (pH 9.5) containing 0.1% SDS. The combined cluates were lyophilized and used to immunize rabbits. Initial injections were administered with complete Freund adjuvant, and booster injections were administered with incomplete adjuvant at 4-week intervals. Antibody titers in sera were monitored in Western blots.

In vitro phosphorylation. Extracellular virions were purified from two 150-cm² culture flasks, washed three times with buffer A (0.1% Nonidet P-40, 20 mM MgCl<sub>2</sub>, 50 mM Tris hydrochloride [pH 8.0]), and collected by centrifugation (30 min, 4°C, 20,000 × g). The protein content was determined in the virus suspension by the Bio-Rad assay. For the kinase reaction 10 μg of protein was incubated with 7 μCi of [γ-32]ATP and 200 μi of kinase buffer (buffer A containing 0.1% Nonidet P-40) for 1 h at room temperature. Then 10% of the reaction was precipitated with trichloroacetic acid, and the incorporation of radioactivity into the protein was determined. Usually about 500,000 cpm/μg of protein was incorporated.

Immunoprecipitation. Immunoprecipitations were performed as described previously (19), with minor modifications. The lysates were precleared by incubation with normal mouse or rabbit serum, and Formalin-fixed Staph-

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ylococcus aureus cells (Behringwerke, Marburg, Focc Republic of Germany) at 4°C for 30 min. The lysates we then clarified by centrifugation at 10,000 × g for 5 min.

RNA extraction and Northern blots. For the preparation of RNA, 70 to 80% confluent monolayers of HFF cells were infected with HCMV strain AD169. Whole cell RNA was extracted from infected ceils as described previously (3). For immediate-early RNA, the cells were maintained in medium containing cycloheximide (50 µg/ml) for 60 mln before infection with 5 to 10 PFU per cell. The cells were infected tor 14 h in medium containing cycloheximide, and RNA was prepared. For early RNA, cells were maintained in medium containing 100 µg of phosphonoacetic acid per ml from 3 to 22 h after infection. For late RNA, cells were harvested and RNA was prepared at 72, 96, and 120 h after infection. The RNA was fractionated on 1.5% agarose-2.2 M formaldehyde gels. Human 18S and 28S rRNA and bacterial 16S and 23S RNAs were used as size markers. RNA was transferred to nitrocellulose membranes as described previously (33):

#### RESULTS

Inotation of a cDNA clone coding for pp28. In our provious studies monospecific antibodies against polypeptides partially purified on preparative polyacrylamide gels were used. The purification of the 28 kDa polypeptide, however, was not feasible by this method since it is not a constituent which is easily identified in preparative gels. We therefore used the monocional antibody P2G11. In Western blots this antibody reacts with a protein of 28 kDa in infected cells and purified virions (22). The antigen recognized by the monocional antibody P2G11 appeared to correspond to the immunogenide 28-kDa polypeptide recognized by immunoglobuling G present in human sera (22).

A total of 150,000 recombinant gt11 phages were sere with P2G11. Two positive signals were obtained. One clone (designated BUML-1) was purified and characterized in detail. E. coll Y1089 cells were infected with the recombinant phages, and the synthesis of the fusion protein was induced by the addition of isopropyl-β-p-thiogalactopy ranoside. In protein extracts of E. coli cells infected with BUML-I an abundant protein of about 130 kDa was synthe sized which was not present in E. coli cells infected with lambda gt11. Here a protein of 118 kDa, corresponding to β-galactosidase, was detectable. Both polypeptides avere readily detectable in polyacrylamide gels after staining with Coomassie brilliant blue, suggesting a high synthesis rate? In Western blots the 130-kDa polypeptide of BUML-1 reacted exclusively with the monocional antibody P2G11 (data not shown). Proteins from lambda gtll-infected cells or fusion proteins from unrelated cDNA clones were not recognized. We took this as evidence that BUML-1 was synthesizing a hybrid protein of B-galactosidase and purts of pp28 of HCMV. To further substantiate this, DNA was prepared from plate lysates and the cDNA insert was cut out with EcoRI, the enzyme used in the original cloning procedure. In agarose gels stained with ethidium bromide a DNA fragment of about 270 bp was detected (date not shown). This fragment was radiolabeled with "P and hybridized in a dot-spot assay to DNA from eight cosmid clones spanning the whole HCMV genome (5). The cosmid pCM1058, containing the fragments HindIII-T, -R, and -E, hybridized to the cDNA In a more detailed Southern blot analysis the complementary sequences were located within a 500-bp KpnI-Smal fragment at the left end of HindIII-R (Fig. 1A). Since HindIII-R is cleaved into tw Kpnl-Smal fragments of equal size [Fig.

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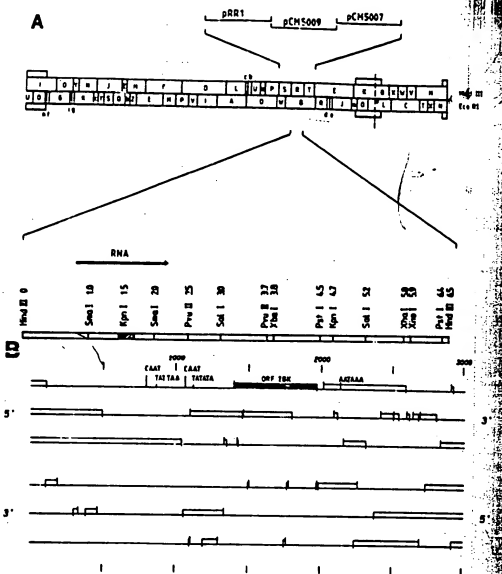


FIG. 1. Structure of the HCMV AD169 genome and localization of the pp28 DNA. (A) Schematic representation of the HCMV genome with the restriction maps for Hindill and EcoRi. The plasmids containing Hindill fragments S (pRR1), R (pCM599), and T (pCM5997) and Indicated on top of the map. The 6.5-kb Hindill R fragment is shown on an expanded scale below the HCMV genome. Only aome relevant sestiletion endonuclease recognition sites are given. The localization of the cDNA of BUML-1 is marked by the hatched area. The orientation of transcription is indicated by an arrow. (B) Reading frame analysis of the first 3,000 nucleotides of Hindill-R (4). Bars above the horizontal line indicate start codons; bars below the lines indicate stop codons. The potential CAT and TATA sites and the poly(A) signal sequence indicated.

1A), it was not possible to determine the exact location of the cDNA complementary sequences in this analysis. However, the two fragments can be distinguished by an Sstll recognition site located at nucleotide 1877. Taking advantage of this restriction site, it was possible to locate the cDNA to the right Kpnl-Smal fragment in the genome rientation shown in Fig. 1A. To establish the nucleotide sequence the 270-bp cDNA fragment was subcloned int M13mpl1 and sequenced by the chain termination method (26). The exact

size of the cDNA was found to be 273 nucleotides (Fig. 2). It contained an open reading frame running through the entire fragment. The translation of 273 bp in addition to the lacz gene would result in a polypeptide of approximately 130 kDa. The size of the fusion protein synthesized in BUML-1 is in good agreement with this theoretical value.

Genomic sequence of the region coding for paid. The nucleotide sequence of a 3-kilobase (kb) segment within Hindill-R was determined by the dideoxy-chain termination

FIG. 2. Nucleotide sequence of the HindIII-R DNA segment encoding pp28. The HindIII site between fragments S and R is taken as zero. For the sake of uniformity with Fig. 1B the sequence has not been renumbered. The cDNA clone BUML-1 (underlined) corresponds to the distribution DNA sequence between nucleotides 1415 and 1688. The TATA consensus sequences and the poly(A) signal sequence AATAAA are boxed. The amino acid sequence of ORF pp28 (Fig. 1B) is given in the one-letter code.

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method. Figure 2 shows the sequence of nucleotides 800 to 2200. The HindIII site between fragments S and R is taken as nucleotide 0. The region of full homology between cDNA and genomic DNA extends from nucleotides 1415 to 1682. This determines the open reading frame (ORF) between nucleotides 1399 and 1967 as the coding region for pp28 if the transcript is not spliced (see below). The AIIC coden at position 1399 follows the known rules for effective translational start sites (12). Analysis of the nucleotide sequence upstream region revealed two potential TATA box requences at nucleotides 893 through 898 and 1133 through 1140. Both are preceded by potential CAAT boxes 83 and 59 mucleotides upstream. However, there is another potential initiation codon that fulfills the rules of K zak between the potential prom ter sequences and the initiation codon in the

pp28 reading frame. This ATG at position 1336 has a purine at position -3 but is followed 7 codons later by a termination codon. A potential polyadenylation signal (AATAAA is located at nucleotides 2144 through 2149. The ORF pp28 is surrounded by two ORFs running in different directions (Fig. 1B). The left ORF starts within HindIII S and extends to nucleotide 1046 (Fig. 2). The right ORF is located at the opposite strand ending at nucleotide 2238. It is worth noting that both left and right ORFs show homology to reading frames BGLF5 and BBRF3 of Epstein-Barr virus. On the Epstein-Barr virus genome (the two reading frames are arranged in a nearly identical fashion (1). N homology to other published herpesylvus sequences was found for pp28 in the numerical molecular size of the polypeptide encoded by ORF pp28 is 20,933 Da. This is a clear discrepancy with the



FIG. 3. Identification of transcripts homologous to RUML-1 in infected cell RNA. Whole cell RNA was isolated under immediate-early (lane 2), early (lane 3), and late (lane 4) conditions and fractionated on a 1.5% agarose gel containing formaldehyde. The gel was blotted onto nitrocellulose and probed with the radiolabeled 473-bp eDNA fragment of BUML-1. Lane 1 contains RNA from mock-infected cells. Each lane co-ains 13 µg of RNA. Fragment is given in kilobases.

size estimated from polyacrylamide gel electrophoresis. The difference between this and the theoretical value of about 7 kDa is most probably due to the phosphorylation (see below) and the unusual characteristics of the protein. Computer analysis by the method of Hopp and Woods (7) revealed that pp28 is an extremely hydrophilic protein. pp28 lacks tryptohan, the most hydrophobic amino acid, and contains only 1% phenylalanine and 1% tyrosine, whereas the hydrophilic residues arginine, aspartic erid, glutamic acid, and lysine make up 33% of the molecule. According to the hydrophilicity pattern the most hydrophilic regions are centered around amino soids 50 through 60 and 110 through 120. The monoclonal antibody P2G11 recognizes an epitope encoded by the cDNA between amino acids 7 and 95. This is onsistent with the prediction of Hopp and Woods that hydrophilic regions are highly antigenic. An overestimation of the molecular size seems to be a phenomenon common to the phosphoproteins of HCMV. Both the basic phosphoprotein and the major matrix protein have apparent molecular sizes of 150 and 65 kDa when estimated by analysis on SDS-polyacrylamide gel electrophoresis. The theoretical values, however, calculated from computer analysis are 113 and 61 kDa, respectively (9, 24).

Analysis of the terminated and time of appearance. In Northern blot anways a the size of the pp28-specific transcript was determined. Total RNA from AD169-infected cells was isolated under immediate-early, early, and late conditions. The RNAs were separated on a 1.5% agarose denaturing gel and probed with the 12P-radiolaheled cDNA BUML-1 fragment. A RNA species of 1.3 kb was the most prominent signal (Fig. 3). This transcript was present only in RNA preparations isolated late in infection. The 1.3-kb RNA was encoded entirely within the HindIII R fragment. In Northern blots with the surrounding fragments Hindliff-T and -S no RNA species of this size could be detected at late times (data not shown). The 1.3-kb RNA was abundant late in the infectious cycle. An additional late RNA of 1.5 kb was also detected within the HindIII R fragment. This RNA is transcribed from coding sequences downstream of ORF pp28, most probably from the right ORF (H. Meyer, M. Mach unpublished). Northern blot analysis with singlestranded cDNA probes confirmed that the direction of transcription for the 1.3-kb RNA was from left to right in the genome arrangement shown in Fig. 1A (data not shown).

Expression cloning of pp28. To express a large part of pp28

PP28 GENE OF HUMAN CYTOMEGALOVIELE

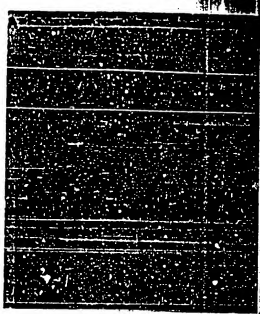


FIG. 4. Recognition of the β-galactosidase-pp28 hybrid profess by human sera. (A and B) E. coll extracts were separated by SDS-polyacrylamide gel electrophoresis on a 8% acrylamide gel. After transfer to nitrocellulose the membrane was cut, and one helf was stained with amido black to control for the transfer (A). This second half was subjected to the immune reaction (B). Lanes (p271) protein attract from clone p271; (pEX2) protein extract from pEX12; (Std) molecular weight marker (Sigma Chemical Co.). Serum (code no. 54527) dilution was 1:750. (C) Proteins from purified virious or dense bodies were separated on a 10% acrylamide gel and subjected to a Western biot analysis with the same human serum samples as int A and B. Lanes: (V) virioms; (Db) dense bodies. Serum (code add 54527) dilution was 1:250. All sizus are given in kilodaltons.

for immunological studies, plasmid p271 was constructed (see Materials and Methods). The vector contains a 500-bp Kpnl-Smal fragment of pp28 fused to the cro-B-galactosides e gene of pEX-2 (29). The Kpnl-Smal fragment codes for 87% of pp28. Fourteen amino acids of the amino terminus and 11 of the carboxy terminus are tacking in clone p271. The synthesis of the hybridprotein is controlled by the P-promoter of phage lambda. Upon induction, a protein of 136 kDa accumulated to high levels in E. coll cells carrying the p271 plasmid (Fig. 4A). This polypeptide was recognized specifically by monoclonal antibody P2G11, indicating that the p271 construct carried the HCMV sequences in the right orientation and reading frame.

pp28-β-galactor dase fasion protein recognized by human sera. To test whether HCMV polypeptides synthesized in Escoli could be used as antigens for diagnostic purposes. Western blots with protein extracts of clone p271 were carried out. In the first set of experiments 14 HCMV-positive human serum samples and 6 negative serum samples were analyzed. Since human sera may contain antibodies against E. coli proteins and especially β-galactosidase, all serum samples were preincubated with an extract of E. coli cells expressing the cro-β-galactosidase protein. The same extracts were also run as controls. An example of a Western blots analysis is shown in Fig. 4. The serum weakly reacted with the pp28 polypeptide when gradient-purified virus was used as the antigen. The reactivity was below detection

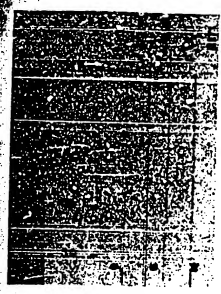


FIG. 5. Detection of pp28 with antisera against recombinant approteins. Proteins from extracellular virious were separated by SDS-polyacrytamide gel electrophoresis on a 10% acrytamide gel. After transfer to nitrocellulose the membranes were cut and subsected to immunereactions with various sera (A). (B) Amount of protein transferred to nitrocellulose, stained with amido black. Lanes: (Vir) extracellular HCMV virious; (M) molecular weight markers; (P2011) monnecional antibudy P2011 dilution 1:250; (p abundle, p a 271) preimmune rabbit sera; (aBUML, a 271) rabbit sera; made against the fusion protein of BUML and p 271, respectively. All interes are given in kilodaltons.

limits when purified dense bodies were used (Fig. 4C). The p271 fusion protein, however, was strongly recognized by the serum even at a fourfold-higher dilution (Fig. 4B). This might be due to the higher amount of antigen present in the p271 protein extract. The preincubation of the human seral with the E. coll extract efficiently eliminated the problems which may arise from anti-E. coll antibodies. Of the 14 human serum samples, 10 recognized pp28 in purified virtions. The same serum samples also reacted with the p271 fusion protein, indicating that a recombinant protein can substitute for the authentic antigen. No reaction was seen with all of the HCMV-negative sera.

An antiserum was also raised against the recombinant protein. Protein extracts from p271 cells were separated on a spreparative 10% polyacrylamide gel, and the 136-kDa hand was cut out after staining with Coomassie brilliant blue. The protein was eluted from the get and used to immunize rabbits. The reactivity of the serum was monitored in Western blots by using purified virions as antigens. After one booster injection the rabbit sera reacted with a polypeptide which showed the same apparent molecular weight as pp28 (Fig. 5). The fast immune response again proves the high immunogenicity of the pp28 protein. Moreover, a serum raised against the BUML fusion protein also recognized the pp28 and did not show any reactivity to other HCMV proteins (Fig. 5). The specificity of the recumbinant sera frepresents additional proof that the identified reading frame within the Hindill R fragment in fact encodes the 28-kDa polypeptide of HCMV. Neither recombinant serum was able to neutralize HCMV strain AD169 grown in tissue culture (data not shown).



FIG. 6. SDS-polyacrylamide gel electrophoretic analysis of immune precipitated proteins from in vitro-phosphorylated HCMV virions. Monoclonal antibody P2G11 (lane 3), a271 rabbit serum (lanes 4 and 5), a rabbit serum against non infectious enveloped particles (lane 1), and a preimmune rabbit serum (lane 2) were used as sources of precipitated antibodies. Precipitated proteins were cluted in the presence of β-mercapto-thanol, analyzed on a 10% acrylamide gel, and exposed to X-tuy film. Lane 6 contains HCMV proteins labeled in vitro with {γ-12PJATP, Lane 7 contains a 1-4CL labeled protein mixture. Molecular sizes (in kilodaltoni) are shown on the right.

Immunoprecipitation of pp28 with different antib Previous studies have described a phosphoprotein (19 Nowak, Ph.D. thesis), a glycoprotein (20), and a capsid protein (8) with molecular weights in the range of 28 kd. To test whether pp28 described in this work was identical with one of these polypeptides, we investigated a potential posttranslational modification of the protein. Digestion of virion proteins with endoglycosidases did not alter the mobility of pp28, suggesting that the protein was not glycosylated to a great extent (data not shown). To analyze a potential phosphorylation of pp28, extracellular HCMV particles were phosphorylated in vitro. HCMV particles have been shown to contain an associated protein kinase which is capable of phosphorylating in vitro the same polypeptides which are phosphorylated in vivo (23). The presence of a virion associated protein kinase is typical of enveloped viruses and has been reported for many members of the herpesvirus group (30, 32). Extracellular HCMV particles were collected from tissue culture supernatants 72 h postinfection. The HCMV particles were separated into virions, noninfections enveloped particles (NIEPS), and dense bodies by a glycere ol-tartrate gradient. The fraction containing virious was collected and used for the in vitro phosphorylation. Five proteins of 150, 71, 65, 36, and 28 kDa were most intensely labeled (Fig. 6). They correspond to the basic phosphopro tein (pp150), upper and lower matrix proteins (pp71 and pp65), assembly protein (pp36), and the 24-kDa phosphoprotein, respectively, as described by Roby and Gibson (23). The presence of the 36-kDa assembly protein indicates that the virus preparation was not completely free of NIEPS. since this polypeptide is an exclusive constituent of NIBPS (2). In agreement with previous studies (23), we also observed that the 71-kDa upper matrix protein was phosphorylated better in vitro when compared with the low

protein of 65 kDa. The in vitro-phosphorylated proteins were immunoprecipitated with monoclonal antibody P2G11 and the rabbit antiserum raised against the p271 fusion protein. Both antibodies precipitated a protein of 28 kDa (Fig. 6). The same polypeptide, was also precipitated, although to a much lesser extent, by a rabbit serum raised against NIEPS, (Fig. 6). In addition the NIEPS antiserum precipitated phosphorylated proteins of 150, 80, 71, 65, 52, and 36 kDa. The reactions are highly specific, since a rabbit preimmune serum did not precipitate any of the phosphorylated proteins. These data strongly suggest that the 28-kDa structural protein of HCMV, which is encoded by the Hindlil R fragment, is a phosphoprotein.

#### DISCUSSION

In this report we describe the identification of the gene cuding for the immunogenic 28-kDa phosphoprotein by using a monoclonal antibody.

After the initial isolation of immunoreactive lambda gt11 HCMV plaques, the genomic region was localized by hybridization analysis with available cosmid and plasmid clones of HCMV DNA. The coding sequence for pp28 is located in the long unique segment of the viral genome within the left end of the Hindill R fragment. In this area three ORFs are located. The nucleotide sequence comparison between cDNA and genomic DNA established that the small ORF codes for pp28. Our data exclude the possibility that pp28 is derived from a spliced transcript containing parts of the left ORF and the small adjacent ORF. The pp28 cDNA spans the gap which would have to be spliced out to create an ORF. The pp28 gene is transcribed into a 1.3-kb mRNA which is present exclusively late in the infectious cycle. In this study we did not attempt to map this RNA precisely. A detailed analysis of the transcripts originating from this part of the Hindlii R fragment will be presented elsewhere (R. Lehner and M. Mach, manuscript in preparation). With the accumulating sequence information on all human herpesviruses it becomes apparent that some regions of the viral genomes show extensive homology. These similarities are reflected in the sequence of the proteins encoded as well as the relative organization of the genes. Examples are the coding regions for the DNA polymerase and the major glycoprotein (1, 6, 11). There are, on the other hand, a number of HCMV genes which do not show homology to those of other herpesylruses. Examples are the 150-kDa basic phosphoprotein, the 65-kDa major matrix protein, and the entire short unique region (35). In pp28 a different pattern seems to exist. An unique gene is located interspersed between reading frames which are conserved between distantly related herpesvi-

Recently, Martinez and St. Jeor (18) described the isolation of a lambda gill clone which also maps within the Hindill R fragment. The fusion protein synthesized by this clone is recognized by human sera. Antibodies raised against the fusion protein recognize a 19-kDa structural polypeptide which is transcribed from a small RNA of about 1.5 kb. Since no precise mapping or sequencing data are given, it is not clear whether both proteins are identical, especially since the Hindill R fragment codes for a number of immunogenic structural proteins (R. Lehner and M. Mach, unpublished data).

In several studies HCMV polypeptides in the range of 28 kDa have been described. In four reports either a posttranslational modification or the localization of the protein within the virus has been investigated. Roby and Gibson (23) as

well as Nowak et al. (19; Nowak, Ph.D. thesis) described phosphoproteins of 24 and 19 kDa, respectively suggested that the proteins are located in the matrix rather than the capsid or the envelope. Pereira et al. (21) described a monoclonal antibody which in immunoprecipitations for acts with a 25-kDa glycosylated protein, a member of the 2D family, in infected cell extracts. Irmiere and Gibson (8) have identified a 28-kDa protein that is present in both the A and B capsids of different HCMV strains. Our results suggest that the 28-kDa protein corresponds to the phosphorylated matrix protein. Consistent with this suggestion are termuno-electron microscopy atudies which localized the protein recognized by the monoclonal antibody P2G11 on the outline of cytoplasmic viral capsids (16).

The fact that pp28 is recognized by the majority of human sera turns it into a candidate for a diagnostic reagent. The most reliable protein in this regard is the basic phosphopro tein (pp150) It is recognized by virtually all sera that we (10) and others (14) have tested and therefore represents a useful thoi to assess the occurrence of a previous HCMV infection Preexisting high anti-pp150 titers, on the other hand may complicate the detection of a recurrent infection if the diagnostic procedure is based exclusively on this protein. Therefore, it seems desirable to include additional anticens in the evaluation of a serological HCMV test. It has been shown that together with pp65 and gp58, antibody titing against p.25 increase sub-tantially during a recurrent HCMV infection (14). Therefore these four proteins (pp150) pp65, gp58, and pp28) might represent an antigenic complex sensitive enough to detect low levels of antihodies due to a past infection and able to detect changes in antibody liters due to a present infection. We have shown here that recombinant protein synthesized in E. coli can substitute for the authentic pp28 antigen. Although the number of sera that we have tested is small, we believe that it is safe to assume that in general this polypeptide produced in E. coll can serve as a reliable antigen for the detection of antibodies against pp28. A study testing this hypothesis on a sufficient number of antisera is currently in progress. To avoid problems which could arise from antibodies against E. coll proteins, it would be advantageous to express pp28 unfused and to purify the protein. These preparations could then be used in enzyme linked immunosorbent assays. So far we have not been able to produce sufficient quantities of pp28 in an unfused state.

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